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Hierarchical Porous Li₂Mg(NH)₂@C Nanowires with Long Cycle Life Towards Stable Hydrogen Storage

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The hierarchical porous Li₂Mg(NH)₂@C nanowires full of micropores, mesopores, and macropores are successfully fabricated via a single-nozzle electrospinning technique combined with *in-situ* reaction between the precursors, i.e., MgCl₂ and LiN₃, under physical restriction upon thermal annealing. The explosive decomposition of LiN₃ well dispersed in the electrospun nanowires during carbothermal treatment induces a highly porous structure, which provides a favourable way for H₂ delivering in and out of Li₂Mg(NH)₂ nanoparticles simultaneously realized by the space-confinement of the porous carbon coating. As a result, the thus-fabricated Li₂Mg(NH)₂@C nanowires present significantly enhanced thermodynamics and kinetics towards hydrogen storage performance, e.g., a complete cycle of H₂ uptake and release with a capacity close to the theoretical value at a temperature as low as 105 °C. This is, to the best of our knowledge, the lowest cycling temperature reported to date. More interestingly, induced by the nanosize effects and space-confinement function of porous carbon coating, a excellently stable regeneration without apparent degradation after 20 de-/re-hydrogenation cycles at a temperature as low as 130 °C was achieved for the as-prepared Li₂Mg(NH)₂@C nanowires.

Solid-state hydrogen storage has been widely considered as the safest and most effective way to achieve the widespread use of H_2 as a prominent energy carrier in the future¹. Despite decades of extensive efforts, it is still a big challenge to find a suitable H-rich material that features high safety, low cost, and fast kinetics for H_2 absorption and desorption under moderate conditions^{2–4}. Metal-N-H systems have been extensively investigated as potential materials for solid-state hydrogen storage since the discovery of the hydrogen storage performance of the Li-N-H system, which was found to reversibly store 11.5 wt. % hydrogen^{5,6}. Inspiringly, it was verified that partial substitution of Mg for Li in the Li-N-H composite to form the Li-Mg-N-H system could significantly decrease the desorption enthalpy change^{7–9}. Among such composites, the Mg(NH₂)₂-2LiH system is one of the most promising hydrogen storage materials due to its favorable thermodynamics (39 kJ mol⁻¹ H₂), which is lowered by 35% compared with the LiNH₂-LiH system, as well as its relatively high H₂ content and good reversibily stored in this system at an operating temperature of 90°C under 1 bar of equilibrium hydrogen pressure, based on the thermodynamic calculation^{12,13}, which could satisfy the practical requirements of proton exchange membrane (PEM) fuel cells.

$$Li_2Mg(NH)_2 + 2H_2 \leftrightarrow Mg(NH_2)_2 + 2LiH$$
(1)

Nevertheless, due to both the sluggish interface reaction between Mg(NH₂)₂ and LiH in the preliminary step and the following mass transport across the thus-formed imide layer, which results in a rather high activation energy barrier (E_a : ~102–120 kJ mol⁻¹), a temperature higher than 200°C is required to achieve a reasonable H₂ sorption rate¹³⁻¹⁵. In addition, another big challenge for practical application of the Li-Mg-N-H system is the undesirable release of ammonia resulting from the insufficient reaction of Mg(NH₂)₂ with LiH, which not only significantly degrades the H₂ cycling capacity due to the loss of elemental N, but also would damage the active catalyst in the PEM fuel cells^{16,17}. Previous experimental results have clearly established that, in terms of mitigating the above-mentioned problems, intimate contact between Mg(NH₂)₂ and LiH through the reduction of their



Figure 1 | Schematic diagram of the in-situ technique to fabricate the carbon-coated Li-Mg-N-H NWs via electrospinning using a single-needle nozzle (top) and typical SEM images of the resulting products at different stages during preparation (bottom).

particle sizes is an effective way of improving the hydrogen storage properties of the Li-Mg-N-H system^{11,18}. It is well-known, however, that particles with small sizes are intrinsically unstable and prone to grow into larger crystallites¹⁹. Thus, during long-term cyclic heat treatment for H₂ release and uptake, phase separation and particle growth are inevitable, which would significantly degrade the sorption kinetics¹⁹. Therefore, an efficient host, which is both capable of uniformly confining the nanostructured morphology of the thus-synthesized complex hydrides during repeated heating and is chemically inert, is essential for preserving the nanosize effects on the active substrates inside.

To this end, nanoconfinement of complex hydrides in porous scaffolds, which can serve as containers, has emerged as an intriguing strategy to synthesize nanostructured materials and therefore improve their hydrogen storage properties¹⁹⁻²⁵. The porous templates can effectively, on the one hand, facilitate the formation of nanoparticles and, on the other hand, maintain the shape and size of the active materials during cycles of hydrogen sorption¹⁹. In general, the preparation process involves melt infiltration or solution infiltration, which requires good wettability of the porous scaffolds by the precursors. Moreover, in order to achieve a relatively high loading level with good distribution, many repetitions of the infiltration steps and the following drying are required²⁶. This makes the nanoconfinement time-consuming and tedious, while not allowing control over the homogeneous distribution of confined composites^{26,27}. More importantly, the Li-Mg-N-H system is composed of multiphase components, and all of them have high chemical activity, low solubility, and/or high melting points, which therefore hinders the direct synthesis of a nanosized Li-Mg-N-H composite by the conventional nanoconfinement technique.

Herein, we report a novel *in-situ* reaction strategy based on an electrospinning technique, which is inexpensive, simple, and versatile, for the preparation of nanowoven materials, consisting of long entangled nanofibers of different materials, to fabricate nanowires (NWs) of Li-Mg-N-H composite uniformly coated with hierarchical porous carbon. As illustrated in Fig. 1, the overall synthetic procedure for the carbon-coated Li-Mg-N-H NWs originates from electrospinning coupled with the *in-situ* fabrication of the porous carbon

coating and the consecutive transformation of the precursors to form the Li-Mg-N-H composite. Due to the special physicochemical properties of the components in the Li-Mg-N-H system, as mentioned above, our synthetic procedure is based on the in-situ reaction between MgCl₂ and LiN₃ as the electrospinning precursors. Upon heat treatment concurrent with carbonization to produce the uniform carbon coating, LiN₃ firstly decomposes into Li₃N with vigorous release of nitrogen, which then reacts with MgCl₂ to produce Mg_3N_2 via a metathesis reaction (2). With further increase of temperature, the interaction between the thus-formed Mg₃N₂ and the residual Li₃N results in the formation of disordered LiMgN-type structures (Fig. S1 and S2 in the Supporting Information)²⁸. Afterwards, annealing under H2 atmosphere induces the hydrogenation according to Equation (3)²⁹, which, as a result, forms the composite of Mg(NH₂)₂ and LiH, with LiCl as the by-product. The formation of Mg(NH₂)₂, LiH, and LiCl, as detected by X-ray diffraction of the bulk precursors after annealing (Fig. S1), could verify the feasibility of this procedure.

$$3MgCl_2 + 2Li_3N \rightarrow Mg_3N_2 + 6LiCl$$
 (2)

$$Mg_{3}N_{2} + 4Li_{3}N + 12H_{2} \rightarrow 3Mg(NH_{2})_{2} + 12LiH$$
 (3)

Results

Scanning electron microscope (SEM) images of the as-electrospun NWs are shown in Fig. 1. When a high voltage is applied between the outer metallic capillary, containing the mixed precursor solution of LiN₃, NH₄MgCl₃, and poly(vinyl alcohol) (PVA), and the collector, the liquid jets are stretched by electrostatic forces to generate textile NWs with a uniform coating of PVA. From the SEM images, the formation of a highly interconnected network of the as-electrospun NWs with a diameter of ~250 nm is clearly observed. Subsequently, the as-collected NWs were dewatered at ~85°C, and the temperature was then programmed to rise to 250°C under dynamic vacuum, meanwhile realizing the transformation of NH₄MgCl₃ into MgCl₂,



Figure 2 | (a) High-magnification SEM image of a single Li-Mg-N-H NW; (b) TEM image of Li-Mg-N-H NWs (inset: high-magnification TEM image (scale bar 5 nm), in which the marked area clearly shows the small and large mesopores coated with carbon). TEM image (c) and the corresponding elemental mapping (d) of the as-prepared Li-Mg-N-H NWs with nanoconfinement of the precursors. (e) High-resolution XRD spectra of the Li-Mg-N-H NWs upon heating to various temperatures.

accompanied by the release of the thus-formed ammonia and hydrogen chloride. Simultaneously, the formation of Li₃N due to the decomposition of LiN₃ was verified by the color change (Fig. S3) to the characteristic color of Li₃N, *i.e.*, ruby red from white, across this heating step. On elevating the calcination temperature to 550°C under N₂ atmosphere, the PVA substrates were then transformed into pyrolysis carbon upon thermal decomposition, which homogeneously puts "carbonaceous clothes" on the active composite inside, accompanied by the metathesis reaction between $MgCl_2$ and Li_3N . As shown in Fig. 1, the carbonized NWs perfectly maintain the structural integrity of the as-electrospun polymers and, compared to polymeric NWs with smooth surfaces, exhibit rough surfaces with small relics and mesoporous structure on the outside wall. This is supposed to stem from the explosive release of N2 during the decomposition of LiN₃, which could push away the surrounding PVA substrate from the point where LiN3 exists, leading to the formation of numerous pores after the carbonization of relatively flexible polymers into relatively stiff carbon sheets at high temperature²⁷. Magnified images of Li-Mg-N-H NWs (Fig. 2a) can further demonstrate the presence of open pores on the external surface, with pore diameters in the range of 2-40 nm. It also was found that the diameter of the as-electrospun PVA NWs was significantly decreased

after carbonization, due to the decomposition of the polymer during calcination³⁰. In the high-resolution transmission electron microscope (HRTEM) image (Fig. 2b), not only the wirelike nanostructure, but also the porous interior with mesopores and macropores can be clearly observed. This verifies the homogenous distribution of pores, which can effectively facilitate the H2 transportation across the carbon matrix, owing to the explosive decomposition of LiN₃ inside the carbonized wires, indirectly confirming the uniform dispersion of Li-Mg-N-H composite. The HRTEM image (inset of Fig. 2b) also reveals the presence of small mesopores (~2.5 nm) inside the disordered carbon sheets. The N2 adsorption-desorption isotherms of the as-prepared NWs exhibit typical type-IV hysteresis, with an obvious increase at a high relative pressure ($P/P_0 = 0.80-0.99$), indicative of the presence of both macropores and mesopores, due to the compression of PVA by the strong power of the N2 stream and the subsequent solidification from carbonization, which agrees well with the TEM results. In addition, the isotherms show increases in adsorption at a low relative pressure, suggesting the presence of a large number of micropores in the NWs after calcination. These pores may be derived from the free volume and microporosity in the carbon NWs that is attributed to the disordered packing of the turbostratic carbon sheets and clusters as a result of the carbonization process of the nanostructured polymers³¹⁻³³. Taking advantage of the one-dimensional (1D) and hierarchical porous architecture, the Li₂Mg(NH)₂@C possesses a relatively large Brunauer-Emmett-Teller (BET) surface area of around 438 m² g⁻¹ and pore volume of 0.86 cm³ g⁻¹ according to the N₂ sorption isotherms (Fig. S4). Moreover, the pore size distribution further demonstrates the formation of mesopores in the as-prepared NWs. Accordingly, on account of the special physico-chemical properties of every individual component in the as-electrospun matrices, the hierarchical porous carbon NWs can be obtained via a simple single-nozzle electrospinning. It is worth noting that the energy dispersive spectroscopy (EDS) elemental map (Fig. 2d) of C coincides well with the structure of the NWs, while the maps of Cl, Mg, and N agree well with the C map, which directly demonstrates the homogeneous dispersion of the precursors inside the carbon NWs.

After the carbothermal treatment and annealing in the H₂ atmosphere, high-resolution X-ray diffraction (HRXRD) verified the formation of LiCl, as shown in Fig. 2e, while other products are undiscernible due to a lack of long-range order. The characteristic peaks assigned to the N-H bonds of Mg(NH₂)₂ can be clearly detected, however, from the Fourier transform infrared (FTIR) spectra (Fig. S5), indicating the presence of $Mg(NH_2)_2$ embedded in the porous NWs. Moreover, the in-situ HRXRD confirms the generation of Li₂Mg(NH)₂ with increasing temperature, providing direct evidence of the successful synthesis of the porous carbon-coated Li₂Mg(NH)₂ NWs. In addition, the surrounding carbon matrix appears to have an amorphous structure, acting as a strong buffer to accommodate the sintering of nanostructured Li2Mg(NH)2 during the cycles of H₂ sorption. As deduced from the Debye-Scherrer equation, the average crystallite size of the carbon-coated $Li_2Mg(NH)_2$ is only ${\sim}4$ nm. Clearly, the electrospinning will, on the one hand, eliminate the tedious process of infiltration and drying and, at the same time, is suitable for good encapsulation and distribution of the precursors inside the as-electrospun NWs³⁴, while the blocking of pores and/or agglomeration of active composite on the outside surface of the templates is usually observed during nanoconfinement, which significantly limits complete pore filling and good dispersion of complex hydrides inside the scaffolds²⁶. Finally, the *in*situ metathesis reaction between MgCl₂ and Li₃N stemming from the well-dispersed precursors inside the thus-formed carbon NWs, followed by the hydrogenation process, leads to the formation of the nanosized Li-Mg-N-H composite with homogeneous distribution. Therefore, as a result of the explosive release of N₂ from the aselectrospun wires upon carbothermal treatment, the obtained car-



Figure 3 | (a) Mass spectra (top) and thermogravimetry curves (bottom) of the carbon-coated Li-Mg-N-H NWs and bulk Mg(NH₂)₂/2LiH composite after complete hydrogenation. Hydrogen desorption (b) and absorption (c) curves of the carbon-coated Li-Mg-N-H NWs at different temperatures, with the ball-milled Mg(NH₂)₂/2LiH composite at 135°C included for comparison. Carbon was not considered as an active component for the hydrogen storage measurements.

bon-coated Li-Mg-N-H-based NWs possess a well-designed 1D nanostructure and a three-dimensional (3D) interconnected texture full of micropores, mesopores, and macropores, which can effectively hinder the growth and agglomeration of the confined nanoparticles during heat treatment and promote H_2 access through the porous carbonaceous framework to interact with the active substrates. This is the first report, to the best of our knowledge, on the *in-situ* synthesis of a carbon-coated Li-Mg-N-H-based system with 3D porous nanostructures.

To compare the dehydrogenation behavior of the Li-Mg-N-Hbased NWs with that of a ball-milled composite of Mg(NH₂)/2LiH, mass spectrometry (MS), thermogravimetry (TG), and volumetric temperature-programmed desorption (TPD) characterizations were conducted, as shown in Fig. 3. The bulk Li-Mg-N-H system after ball milling starts to dehydrogenate above 140°C, with dehydrogenation peaking at \sim 199°C, along with the simultaneous emission of ammonia, which is generally consistent with reports in the literature³⁵⁻⁴¹. In comparison, the carbon-coated Li-Mg-N-H NWs exhibited onset and peak temperatures of H₂ desorption that were downshifted to ~78°C and 113°C, respectively, which are 62°C and 86°C lower, respectively, than for their ball-milled counterpart. It is also noteworthy that the generation of the toxic by-product, *i.e.*, NH₃, was significantly depressed in the thermal dehydrogenation of the Li-Mg-N-H NWs, which is favorable for both reversibility and onboard practical applications as a hydrogen-storage medium. The quantitative H₂ desorption measurement by the TG method confirmed the dramatically reduced operating temperature of the carbon-coated Li-Mg-N-H NWs, which corresponds well with the MS spectra. A weight loss of 5.5 wt. % H₂ could be achieved in the temperature range between 78°C and 155°C. From the in-situ HRXRD results (Fig. 2e), the formation of Li₂Mg(NH)₂ upon dehydrogenation is confirmed to start at a temperature of $\sim 110^{\circ}$ C, which is consistent with the TG-MS results.

The isothermal H_2 desorption and absorption properties (Fig. 3b, c) were further characterized by volumetric measurements to clarify the effects of both the reduction in the particle size and the carbonaceous coating with highly porous nanostructure on the modification of the Li-Mg-N-H system. The ball-milled composite shows a H_2 capacity of approximately 0.5 wt. % and 0.65 wt. % for desorption

and absorption, respectively, within 60 min, even at the high temperature of 135°C. On the contrary, the carbon-coated Li-Mg-N-H NWs display a fast hydrogenation at 135°C, with a capacity reaching up to 5.5 wt. %, and a subsequent hydrogen release as high as \sim 5.2 wt. % can be implemented over the period of 60 min. In particular, upon decreasing the operating temperature to 105° C, the H₂ sorption from the Li-Mg-N-H NWs can still proceed, and a complete cycle of H₂ uptake and release (\sim 5.5 wt. %) can be achieved with an extended holding time (Fig. S6). In order to quantitatively estimate the enormously enhanced hydriding/dehydriding kinetics for the asprepared carbon-coated Li-Mg-N-H NWs, the apparent activation energies (E_a) during the H₂ absorption/desorption were determined by combining the Johnson-Mehl-Avrami equation with the Arrhenius equation after fitting the experimental results at various temperatures. Based on the slope of the linear plots of $\ln(k)$ (k, rate constant) versus 1/T(T, absolute temperature), the activation energy for hydrogenation and dehydrogenation of the porous carboncoated Li-Mg-N-H NWs was verified to be around 23.9 and 24.5 kJ mol⁻¹ H₂, respectively, very much lower than for the bulk counterpart (Fig. S7). This directly confirms the significant enhancement of kinetics for the carbon-coated Li-Mg-N-H NWs, due to the reduction of particle size and therefore diffusion distances.

Another important issue with Li-Mg-N-H composite for hydrogen storage application is the significant degradation of H₂ capacity upon cycling, owing to the release of ammonia, and the particle growth and phase separation during thermal treatment. The cycling behaviour of the carbon-coated Li-Mg-N-H NWs, in comparison with their bulk counterpart, is shown in Fig. 4 and Fig. S8, using isothermal dehydrogenation and hydrogenation. The H₂ capacity, as cycling of the bulk Li-Mg-N-H composite proceeds, is severely degraded from 0.7 wt. % to 0.35 wt. % after only 5 cycles of sorption, which is equal to a capacity retention lower than 50%, even with a dwell time of 400 min for one cycle at 130°C. In contrast, in the case of the carbon-coated Li-Mg-N-H NWs cycled under the same conditions, but over a significantly shorter time, the H₂ capacity still reaches 5.3 wt. % and the capacity retention is up to 94.6% after 20 consecutive cycles of hydrogenation and dehydrogenation. The characteristic peaks of Mg(NH₂)₂ and Li₂Mg(NH)₂ can be clearly observed, even through 20 cycles of sorption (Fig. S9), and the porous structure of the Li2Mg(NH)2@C NWs is well maintained after cyclic heat treatment (Fig. S10). Furthermore, the curves of absorption and desorption during the cyclic H₂ sorption are almost identical to each other (Fig. S11), indicating the greatly enhanced hydrogen storage performance, *i.e.*, capacity and kinetics were well-preserved, which is attributed to a strong tolerance of the porous carbonaceous coating towards particle growth and sintering.

Discussion

It is well known that the high activation energy barriers for both the hydrogenation and the dehydrogenation of the Li-Mg-N-H composite results from the sluggish interface reaction due to the slow diffusion of atoms (e.g., Li^+ , Mg^{2+} , and H^+ ions) across amide-imide and imide-hydride phase boundaries and the mass transport along the as-formed layer of imides^{18,42}. The reduction of particle size into nanometer range can significantly decrease the diffusion pathways across amides/imides and hydrides, increase their surface area and develop closer proximity between Mg(NH₂)₂ and LiH, which would lead to the significantly improved kinetics and thermodynamics towards superior hydrogen storage performance. In the case of the ball-milled Li-Mg-N-H composite, the sizes of most of the particles are larger than 1 μ m (Fig. S12), and therefore, the surface-to-volume ratio is small, which endows it with high activation energy. By contrast, it is notable that the *in-situ* fabrication of the carbon-coated Li₂Mg(NH)₂ via electrospinning followed by calcination and annealing can, on the one hand, realize the direct synthesis of nanosized $Li_2Mg(NH)_2$ (~4 nm), significantly improving its hydrogen storage





Figure 4 | (a) Reversible H_2 sorption performance of the carbon-coated Li-Mg-N-H NWs at 130°C, and (b) H_2 capacity dependence of the carbon-coated Li-Mg-N-H NWs (green) and their ball-milled counterpart (black) on cycle number.

performance (Fig. 3), while, on the other hand, the space-confinement function of the porous carbon coating can well preserve the nanostructural features down to the nanometer scale during consecutive cycles of heating, which therefore gives the $Li_2Mg(NH)_2$ stable reversibility without apparent degradation, even up to 20 cycles of de-/re-hydrogenation (Fig. 4).

In summary, we have successfully prepared hierarchically porous Li₂Mg(NH)₂@C NWs via a simple and scalable single-nozzle electrospinning technique with a subsequent carbothermal reaction, which realizes the *in-situ* formation of a porous carbon coating serving as template, with the Li-Mg-N-H composite produced via the subsequent reaction under physical restriction. Our results obviously verified the significantly improved hydrogen storage performance, including complete reversibility at a temperature as low as 105°C and tremendous reduction of the hydriding/dehydriding temperature of the Li₂Mg(NH)₂, with the reduction of particle size down to the nanometer scale. Induced by the space-confinement of the porous carbon coating, the Li₂Mg(NH)₂@C NWs present very good cycling stability, close to the theoretical value over 20 cycles of de-/re-hydrogenation at 130°C. To the best of our knowledge, no such highly stable cycling at relatively low temperature has been reported for Li-Mg-N-H based materials. However, it should be noted that, due to the extra weight of the porous carbon NWs and the by-product of LiCl, the actual reversible H₂ capacity in the whole system is limited. Therefore, further works in the search of an effective solvent to remove the inactive LiCl and/or a new synthetic strategy to achieve a high loading percent of precursors are still required to achieve a high hydrogen storage capacity in the whole system. Nonetheless, these results demonstrate the potential of this strategy in further improving the hydrogen storage performance of complex hydrides and designing the nanowire-like structure with a wide range of materials for developing high-performance energy storage applications.

Methods

Synthesis of hierarchical porous Li2Mg(NH)2@C nanowires. Poly(vinyl alcohol) (PVA, 0.5 g, Alfa Aesar, $M_v = 88000$, CAS: 9002-89-5) was mixed with deionized water (5 mL) and stirred in a water bath at 90°C for 8 h to make the PVA solution. After cooling down to room temperature, LiN3 solution (0.5 mL, 20 wt. %) and NH4MgCl3 (0.05 g) in water were then added and vigorously stirred for 6 h to make a homogeneous spinning dope. The resultant precursor solution was poured into a syringe with a 18-gauge blunt tip needle. The flow rate of the solution was approximately 500 $\mu L~h^{-1}$ and was controlled by a syringe pump (Longer, TJP-3A, China). A grounded stainless steel plate was horizontally placed 10 cm from the needle to collect the as-electrospun NWs. A high voltage of 15 kV was applied by a high-voltage power supply. The as-collected electrospun nanowires were firstly dewatered at a temperature of 85°C under dynamic vacuum for 15 h and then calcined at 550°C for 3 h under dynamic N2 atmosphere to obtain the porous carbon NWs containing the precursors (heating rate 1°C min⁻¹). Finally, the thus-formed carbon NWs were annealed under 150 bar H_2 at 300°C for 12 h for a complete hydrogenation, as schematically illustrated in Fig. 1, and then further annealed under vacuum for dehydrogenation at 150°C, which leads to the formation of the hierarchically porous Li2Mg(NH)2@C NWs. According to elemental analysis, the resulting nanowires contained 9.86 mass% Li, 5.7 mass% Mg, 6.23 mass% N, 2.8 mass% O, 16.7 mass% Cl, and 58.5 mass% C, which gives the mass concentration of Li₂Mg(NH)₂ of ~15.2 mass% in the as-prepared carbon NWs.

Preparation of the ball-milled composite. The ball-milling of Mg(NH₂)₂ and LiH with a molar ratio of 1 : 2 was conducted via a planetary QM-1SP2 for 10 h. The ball-to-powder ratio was 30: 1, with a milling speed of 500 rpm. The milling procedure was carried out by alternating between 30 min of milling and 6 min of rest. In order to prevent contamination by air, all handling and manipulation of the materials were performed in an argon-filled glove box with a recirculation system to keep H₂O and O₂ levels below 1 ppm.

Materials characterization. Mass spectrometry (MS; Hidden HPR 20) was conducted from room temperature, using a heating rate of 2° C min⁻¹ under dynamic nitrogen with a purge rate of 80 ml min⁻¹. Differential scanning calorimetry (DSC) measurements were performed with a TAQ 2000 DSC under N₂ with a gas flow of 40 mL min⁻¹ at a heating rate of 2° C min⁻¹. Nitrogen absorption/desorption isotherms (Brunauer-Emmett-Teller (BET) technique) at the temperature of liquid nitrogen via a Quantachrome NOVA 4200e instrument were collected to characterize the pore structure of the samples. The pore volumes and pore size distributions were obtained by using the Barrett-Joyner-Halenda (BJH) model from the adsorption branches of isotherms. The phase composition of the powders was analyzed by X-ray diffraction (XRD, D8 Advance, Bruker AXS) with Cu K\alpha radiation. Amorphous tape was used to prevent any possible reactions between the sample and air during the XRD measurement. High-resolution X-ray diffraction data were collected by a Mythen-II detector at the Powder Diffraction Beamline, Australian Synchrotron. The sample was loaded into a pre-dried 0.7 mm boron-silica glass capillary tube, and the capillary tube was then sealed with vacuum grease in a glove box filled with argon. In order to identify the phase transformations during dehydrogenation, time-resolved in-situ measurements were conducted using a Cyberstar hot-air blower to heat the capillary from room temperature to the desired temperature, at a constant heating rate of 2°C min-1. Fourier transform infrared (FTIR, Magna-IR 550 II, Nicolet) analysis was conducted to determine the chemical bonding. During the FTIR measurements (KBr pellets), samples were loaded into a closed tube with KBr for measurement in an argon-filled glove box. The morphology of the samples was evaluated using a field emission scanning electron microscope (FE-SEM, JEOL 7500FA, Tokyo, Japan) and a transmission electron microscope (TEM, JEOL 2011 F, Tokyo, Japan). Elemental analysis was performed with an Elemen Tar Vario EL3 Elemental Analyser.

The hydrogen storage properties of the thus-synthesized Li₂Mg(NH)₂@C NWs were investigated on a Sievert's apparatus, denoted as a gas reaction controller (GRC, Advanced Materials Corp., USA). The apparatus was carefully calibrated from the H₂ sorption of a LaNi₅ reference sample with an accuracy of \pm 1%, and, typically, a \sim 800 mg sample was loaded into a stainless-steel autoclave for hydrogenation and dehydrogenation measurements. The H₂ absorption kinetics measurements were performed at various temperatures with an initial pressure of 35 atm, and the desorption properties were detected at various temperatures under a hydrogen pressure below 0.02 atm. The pressure-concentration isotherm (PCI) measurements was 600 s. For comparison purposes, the carbon scaffold masses were excluded when determining the amount of hydrogen released from the relevant composites containing carbon scaffolds.

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Author contribution

G.L.X., Z.P.G., H.K.L. and X.B.Y. proposed, planned, and designed the project. G.L.X., Y.B.T., D.L. and Z.W.L. performed the material preparation, characterizations, and hydrogen storage tests. All authors contributed to writing the manuscript.

Additional information

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